PCT

WORLD INTELLIBETUAL PROPERTY ORGANIZATION International Bureau



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classic	Scation 5:		(11) International Publication Number:	WO 94/18968
A61K 31/415		Ài	(43) International Publication Date: 1 Sep	explor 1994 (01.09.94)
(21) International Application N	18 February 1994			(B1) Designated States: All. CA. IP. Em. ČH, DE, DK, ES, FR, GB, GR, IE, SE).	
(36) Priority Data: 08/018,835 18	February 1993 (18.02.5	73) 1	u s	Published With insernational search report.	
(71) Applicant: PRESIDENT A COLLEGE [US/US]; 124 MA 02138 (US).	ND FELLOWS OF I	IARVAI Cambrid	20 gr.		
(72) Inventors: HALPERIN, Jos MA 02146 (US). BRUGN Newton Highlands, MA (VARA, Carlo; 33 Aberd	Brooklin eene Stre	DE, XCI,		
(74) Agent: GATES, Edward, R Atlantic Avenue, Boston,	.; Wolf, Greenfield & MA 02210 (US).	Sacks, 6	i00	AND WOMATIONS	
				PO SOX CONTROL OF CONTROL OF	94976-0405
				(415) 927-0340 • FAX (415) 9	27-7250
			- 1		
(54) Title: METHODS FOR TR	EATING ARTERIOSO	LEROS	ıs		
(57) Abstract					
(57) Abstract	i a particular class of i	midazole	s du	t inhibit endothelial cell, vascular smooth m	uscle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	t inhibit endothelial cell, vascular smooth m iety of arteriosclerotic conditions.	uscle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	at inhibit endothelial cell, vascular smooth me ety of arteriosclerotic conditions.	ascle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	t inhibit endothelial cell, vascular smooth miety of arteriosclerotic conditions.	ascle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	it inhibit endothelial cell, vascular smooth m iety of arteriosclerotic conditions.	uscle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	it inhibit endothelial cell, vascular smooth m ety of arteriosclerotic conditions.	uscle cell and fibroblasi
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	it inhibit endothetial cell, vascular smooth m ety of arteriosclerotic conditions.	uscle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	at inhibit endothelial cell, vascular smooth meety of arteriosclerotic conditions.	ascle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	t inhibit endothetial cell, vascular smooth m ety of arteriosclerotic conditions.	ascle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	at inhibit endothelial cell, vascular smooth moiety of arteriosclerotic conditions.	ascle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	at inhibit endothelial cell, vascular smooth meety of arteriosclerotic conditions.	ascie cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	it inhibit endothetial cell, vascular smooth meety of arteriosclerotic conditions.	ascle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	t inhibit endothelial cell, vascular smooth metry of arteriosclerotic conditions.	ascle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	it inhibit endothetial cell, vascular smooth meety of arteriosclerotic conditions.	uscle cell and fibroblas
(57) Abstract The applicant has identified	i a particular class of i	midazole	s du	t inhibit endothelial cell, vascular smooth metry of arteriosclerotic conditions.	ascle cell and fibroblas

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Ametria	G₿	United Kingdom	34CR	Marritania
AU	Austrație	GE	Georgia	MW	Maleri
113	Berbados	GN	Cidates	NE	Mar
16	Belgiana	GR	Greace	NL	Netherlands
u	Buckina Pago	HU	Hangury	NO	Marway
BG	Bulgida	R	Ireland	NZ	New Zealand
ม	Benig	π	Italy	PL.	Potenti
ir	Brazil	IP.	Japan	PT	Portugal
BY	Belocus	E E	Kenya	RO	Romania
CA	Canada	KG	Kyrgystan	RU	Reseton Pederation
or or	Cantral African Republic	D	Democratic People's Republic	50	Sudma
CG CG	Congo	_	of Korea	575	Sweden
CE	Switzeriend	13	Republic of Korea	SI	Slovenia
a	Cite d'Ivoire	KZ.	Kazakhetan	SIK	Slovakia
CM	Cameroon	<u>a</u>	"' Dischtenstein	SN""	Seciogal "
CN CN	Calma	ÜK	Sri Lanka	TD	Ched
CS CS	Czechoslovalda	ᄧ	Luxembourg	TG	Togo
CZ.	Czech Republic	ĹŸ	Latria	π	Tajikkan
DE	Germany	MC	Monaco	π	Trinidad and Tobago
DK	Desmark	MD	Republic of Moldova	UA	Ukraine
ES	Spain	MG	Madagascur	US	United States of America
n	Paleod	ML	Mall	UZ	Uphakistan
73	Prance	MN	Mongolia	VN	Vist Nam
	Gabon				
GA	CHIOUG				

METHODS FOR TREATING APTERIOSCLEROSIS

Field of the Invention

The invention relates in general to the field of arteriosclerosis and more particularly to the use of imidazoles that inhibit the Ca⁺⁺ activated potassium channel in arresting endothelial cell, smooth muscle cell and fibroblast proliferation.

Background of the Invention

Arteriosclerosis is a term used to describe a thickening and hardening of the arterial wall. It is believed to be responsible for the majority of deaths in the United States and in most westernized societies. Atherosclerosis is one type of arteriosclerosis that is believed to be the cause of most coronary artery disease, aortic aneurysm and arterial disease of the lower extremities, as well as contributing to cerebrovascular disease. Atherosclerosis is the leading cause of death in the United States.

A normal artery typically is lined on its inner-side only by a single layer of endothelial cells, the intima. The intima overlays the media, which contains only a single cell type, the smooth muscle cell. The outer-most layer of the artery is the adventitia. With aging, there is a continuous increase in the thickness of the intima, believed to result in part from migration and proliferation of smooth muscle cells from the media. A similar increase in the thickness of the intima also occurs as a result of various traumatic events or interventions, such as occurs when a balloon dilatation procedure causes injury to the vessel wall. To date, there is no proven treatment for atherosclerosis.

Imidazoles are synthetic antifungal agents that are used both topically and systemically. Indications for their use include ringworm, tinea versicolor and mucocutaneous candidiasis. These compounds are believed to act by

inhibiting ergosterol synthesis in the fungal c 11 wall, and when giv n topically may cause direct damage to the cytoplasmic membrane.

The fungi comprise five widely differing classes of primitive flora, and the variation in cell physiology and biochemistry are extreme. As a result, most antifungal agents have a very narrow spectrum of antifungal activity.

Various imidazoles have been suggested as treatments for prostate cancer. The only one known to the applicants to have been tested, ketoconazole, appears to inhibit, in high doses, testicular and adrenal synthesis of steroid hormones, including testosterone. The ability of ketoconazole to block steroid synthesis is effective in treating some prostate cancers because proliferation of certain prostate cancer cells is highly dependent upon testosterone. Thus, ketoconazole has been used as a hormonal adjuvant for prostate cancer treatment; it reduces plasma testosterone to castration levels. Ketoconazole, as will be described below, is not useful for inhibiting endothelial or smooth muscle cell proliferation.

Summary of the Invention

The applicants have identified a particular class of imidazoles that inhibit endothelial and/or vascular smooth muscle cell proliferation. The applicants also have identified imidazoles that inhibit proliferation of fibroblasts. These imidazoles can be used to beneficially treat a variety of arteriosclerotic conditions and other conditions characterized by complications involving fibrosis, as described below.

According to one aspect of the invention, a method for treating an arteriosclerotic condition is provided. An imidazole is administered to a subject in need of such treatment. The imidazole is an inhibitor of the Ca⁺⁺ activated potassium channel and is an inhibitor of endothelial and vascular smooth muscle cell proliferation.

The m thods are particularly useful for subjects who have sustained an injury to a blood vessel. Preferred imidazoles are clotrimazole, miconazole and econazole.

According to another aspect of the invention, a method for inhibiting endothelial and vascular smooth muscle cell proliferation is provided. It involves the treatment of endothelial cells or vascular smooth muscle cells of a species with the imidazoles described above. The imidazoles inhibit the Ca⁺⁺ activated potassium channel of erythrocytes of the species. Such methods may be in vivo or ex vivo.

Likewise, methods for treating medical conditions involving fibrosis and for inhibiting the growth of fibroblasts are provided. Such methods involve contacting cells, tissues or subjects with imidazoles that inhibit the Ca⁺⁺ activated potassium channel.

Preferably the cells are in a preparation or in a tissue that is substantially free of fungi. As such, the treatment typically is for cells, tissues or subjects that are otherwise free of indications for the preferred imidazoles.

Brief Description of the Drawings

Figure 1 is a graph illustrating the ability of clotrimazole to inhibit cell proliferation in vascular smooth muscle cells, and the ability to reverse the effects of clotrimazole treatment.

Figure 2 is a graph showing that clotrimazole inhibits DNA synthesis in a dose-dependent fashion.

Figure 3 is a graph comparing the effect upon cell proliferation of a variety of drugs.

Figure 4 is a graph comparing the effect upon the Ca⁺⁺ activated potassium channel of the same drugs tested in connection with Figure 3.

Figure 5 is a graph illustrating the inhibitory effect that clotrimazole has upon complement-induced release of mitogenic activity from endothelial cells.

Detailed Description of the Pr f rred Embodiments

The invention is used in connection with treating arteriosclerotic conditions. An arteriosclerotic condition as used herein means classical atherosclerosis, accelerated atherosclerosis, atherosclerosis lesions and any other arteriosclerotic conditions characterized by undesirable endothelial and/or vascular smooth muscle cell proliferation, including vascular complications of diabetes.

Proliferation of vascular smooth muscle cells is a main pathological feature in classical atherosclerosis. Liberation of growth factors from endothelial cells, it is believed, stimulates the proliferation of subintimal smooth muscle which, in turn, reduces the caliber and finally obstructs the artery. The invention is useful in inhibiting such proliferation and, therefore, in delaying the onset of, inhibiting the progression of or even halting the progression of such proliferation and the associated atherosclerotic condition.

Proliferation of vascular smooth muscle cells produces accelerated atherosclerosis which is the main reason for failure of heart transplants that are not rejected. This proliferation also is believed to be mediated by growth factors and can result ultimately in obstruction of the coronary arteries. The invention is useful in inhibiting such obstruction and reducing the risk of or even preventing-such-failures.

Vascular injury also can result in endothelial and vascular smooth muscle cell proliferation. The injury can be caused by any number of traumatic events, or interventions, including vascular surgery and angioplasty procedures performed for example by balloon dilatation catheters. Re-stenosis is the main complication of successful balloon angioplasties of the coronary arteries. It is believed to be caused by the release of growth factors as a result of mechanically injuring the endothelial cells lining the

WO 94/18968

PCT/US94/01749

-5-

coronary art ries. The invention can be useful in inhibiting unwanted endothelial and smooth muscle cell proliferation and delaying or even avoiding altogether re-stenosis.

Other arteriosclerotic conditions include diseases of the arterial wall that include proliferation of endothelial and/or vascular smooth muscle cells, such as vascular complications of diabetes, diabetic glomerulosclerosis and diabetes retinopathy.

Other uses of the invention include by-pass surgery, coronary by-pass surgery, and procedures in addition to balloon angioplasty for re-establishing patency in occluded or partly occluded vessels, e.g. atherectomy, laser procedures and ultrasonic procedures.

The invention also is useful in treating fibrosis and other medical complications of fibrosis, all resulting in whole or in part from the proliferation of fibroblasts.

Medical conditions other than atherosclerosis include undesirable tissue adhesion resulting from surgery or injury.

The invention is used in connection with treating subjects having, suspected of having, developing or suspected developing such conditions.

A subject as used herein means humans, primates, horses, cows, pigs, sheep, goats, dogs, cats and rodents.

The imidazoles useful in the invention also inhibit endothelial and/or vascular smooth muscle cell proliferation. Imidazoles useful in the invention also inhibit the proliferation of fibroblasts. Inhibition of such

It was not expected that inhibitors of the Ca⁺⁺ activated potassium channel would inhibit endothelial cell, vascular smooth muscle cell, or fibroblast proliferation. Other specific inhibitors of the Ca⁺⁺ activated potassium channel (such as charybdotoxin, caliotoxin and iberotoxin) do not inhibit proliferation of endothelial or vascular smooth muscle cells. Moreover, inhibitors of other transport systems that are activated by mitogens, such as ouabain (highly specific inhibitor of the Na/K pump) and amiloride (inhibitor of Na/H exchange) do not inhibit cell proliferation. Thus, the results obtained by the inventor are surprising.

Without limiting the invention to the use of the specific compounds listed, the following is a list of preferred compounds and salts thereof useful in the methods of the invention.

Clotrimazole

IH-Imidazole, 1-[(2-chlorophenyl)diphenylmethyl]-.
---Lotrimin (Schering); Mycelex (Miles)-----



1-(o-Chloro-α,α-diphenylbenzyl)imidszole [23593-75-1] C22H₁₇CIN₂ (344.84).

Econazol

14.1miderale (+).1.12.1(4-chlorophenyl)methovyl-2-(2.4-chlorophenyl)ethyl]-, mononitrate, Ecostatin (Squibb)

(±)-1-[2,4-Dichloro- β -[(p-chlorobenzyl)oxy]phenethyl]imidazole moņopitrate [68797-31-9] $C_{18}H_{15}Cl_3N_2O$.

Econazole Nitrate

1H-Imidazole, (±)-1-[2-[(4-chlorophenyl)methoxy]-2-(2,4-... dichlorophenyl)ethyl]-, mononitrate, Ecostatin (Squibb)

 $\label{eq:continuous} $$(\pm)-1-[2,4-Dichloro-$\beta-[(p-chlorobenzyl)oxy]$ phenethyl]imidazole mononitrate [68797-31-9] $C_{18}H_{15}Cl_3N_7O.HNO_3$ (440.70).$

Miconazole

1H-Imidazole, 1-[2-(2.4-dichlorophenyl)-2-[(2,4-dichlorophenyl)methoxy[ethyl]-, Monistat (Janssen)

.....1-[2,4-Dichloro-β-[(2,4-dichlorobenzyl)oxy|phenethyl]imidazole-[22916-47-8] C₁₈H₁₄Cl₄N₂O (416.12).

Miconazole Nitrate

Monistat (Ortho)

[22832-87-7] $C_{18}H_{14}Cl_4N_2O.HNO_3$ (479.15).

The above imidazoles are well recognized, pharmacologically characterized, and licensed for use by the FDA today either as antimycotic agents or antiprotozoal

agents. As such, established and mpirically documented paremeters regarding their limited toxicity and their useful dosages are well described in the scientific and medical literature.

The imidazole used in the methods of the invention may be administered per se (neat) or in the form of a. pharmaceutically acceptable salt. When used in medicine, the salts should be both pharmacologically and pharmaceutically acceptable, but non-pharmaceutically acceptable salts may conveniently be used to prepare the free active compound or pharmaceutically acceptable salts thereof. Pharmacologically and pharmaceutically acceptable salts include, but are not limited to, those prepared from the following acids: hydrochloric, hydrobromic, sulphuric, nitric, phosphoric, maleic, acetic, salicyclic, p-toluenesulfonic, tartaric, citric, methanesulphonic, formic, malonic, succinic, naphthalene-2-sulphonic, and benzenesulphonic. Also, pharmaceutically acceptable salts can be prepared as alkaline metal or alkaline earth salts, such as sodium, potassium or calcium salts of the carboxylic acid group. Thus, the present invention involves the use of pharmaceutical formulations which comprise certain imidazoles together with one or more pharmaceutically acceptable carriers and optionally other therapeutic ingredients. The carrier(s) and other ingredients of course must be pharmaceutically acceptable.

Analogs of the foregoing compounds that act as functional equivalents also are intended to be embraced as equivalents and within the scope of the invention.

A variety of administration routes are available. The particular mode selected will depend, of course, upon the particular drug selected, the particular condition being treated and the dosage required for therapeutic efficacy. The methods of this invention, generally speaking, may be practiced using any mode of administration that is medically acceptable, meaning any mode that produces therapeutic levels

of th imidazoles f the invention without causing clinically unacceptable adverse effects. Such modes of administration include oral, rectal, topical, nasal, transdermal or parenteral (e.g. subcutaneous, intramuscular and intravenous) routes. Formulations for oral administration include discrete units such as capsules, tablets, lozenges and the like. Other routes include intrathecal administration directly into spinal fluid, direct introduction onto an arterial surface such as by various catheter and balloon angioplasty devices well known to those of ordinary skill in the art, and intraparenchymal injection into targeted areas on an organ such as a heart.

The compositions may conveniently be presented in unit dosage form and may be prepared by any of the methods well known in the art of pharmacy. Such methods include the step of bringing the active imidazole into association with a carrier which constitutes one or more accessory ingredients. In general, the compositions are prepared by uniformly and intimately bringing the imidazole into association with a liquid carrier, a finely divided solid carrier, or both, and then, if necessary, shaping the product.

Compositions of the present invention suitable for oral administration may be presented as discrete units such as capsules, cachets, tablets or lozenges, each containing a predetermined amount of the imidazole, in liposomes or as a suspension in an aqueous liquor or non-aqueous liquid such as a syrup, an elixir, or an emulsion.

Compositions suitable for parenteral administration conveniently comprise a sterile aqueous preparation of the imidazole, which is preferably isotonic with the blood of the recipient. This aqueous preparation may be formulated according to known methods using those suitable dispersing or wetting agents and suspending agents. The sterile injectable preparation may also be a sterile injectable solution or suspension in a non-toxic parenterally-acceptable diluent or solvent, for example as a solution in polythylene glycol and

lactic acid. Among the acceptable v hicles and solv nts that may be employed ar water, Ring r's solution and isotonic sodium chloride solution. In addition, sterile, fixed oils are conventionally employed as a solvent or suspending medium. For this purpose any bland fixed oil may be employed including synthetic mono- or diglycerides. In addition, fatty acids such as oleic acid find use in the preparation of injectibles.

Other delivery systems can include sustained release delivery systems. Preferred sustained release delivery systems are those which can provide for release of the imidazoles of the invention in sustained release pellets or capsules. Many types of sustained release delivery systems are available. These include, but are not limited to: (a) erosional systems in which the imidazole is contained in a form within a matrix, found in U.S. Patent Nos. 4,452,775 (Kent), 4,667,014 (Nestor et al.); and 4,748,024 and 5,239,660 (Leonard) and (b) diffusional systems in which an active component permeates at a controlled rate through a polymer, found in U.S. Patent Nos. 3,832,252 (Higuchi et al.) and 3,854,480 (Zaffaroni). In addition, a pump-based hardware delivery system can be used, some of which are adapted for implantation.

Oral administration for many arteriosclerotic conditions will be preferred because of the convenience to the patient, although topical and localized sustained delivery-may-be even-more desirable for certain treatment regimens.

The imidazoles, when used in vivo, are administered in therapeutically effective amounts. A therapeutically effective amount means that amount necessary to delay the onset of, inhibit the progression of or halt altogether the onset or progression of the particular condition being treated. Such amounts will depend, of course, on the particular condition being treated, the severity of the condition, and individual patient parameters including age, physical condition, size, weight and concurrent treatment.

Thes factors ar will known to those of ordinary skill in the art and can be addressed with no more than routine experimentation. It is preferred generally that a maximum dose be used, that is, the highest safe does according to sound medical judgment. It will be understood by those of ordinary skill in the art, however, that a patient may insist upon a lower dose or tolerable dose for medical reasons, psychological reasons or for virtually any other reasons.

Generally, daily oral doses of active compound will be from about 0.01 milligrams/kg per day to 1000 milligrams/kg per day. Small does (0.01 - 1 mg) may be administered initially, followed by increasing doses up to about 1000 mg/kg per day. In the event that the anti-arteriosclerotic response in a subject is insufficient at such doses, even higher doses (or effective higher doses by a different, more localized delivery route) may be employed to the extent patient tolerance permits. Multiple doses per day are contemplated to achieve appropriate systemic levels of compounds.

EXAMPLES

Materials

Abbreviations: ChTX, Charybdotoxin; CLT, clotrimazole; ECZ, econazole; MCZ, miconazole; FCZ, fluoconazole; METZ, metronidazole; IbTX, iberotoxin; KTX, kaliotoxin; DIDS, di-isothiocyano-disulfonyl stilbene; hemoglobin concentration; MCHC, mean corpuscular hemoglobin concentration; MOPS, 3-[N-morpholino]propanesulfonic acid.

Drugs and Chemicals

Synthetic charybdotoxin (ChTX) was purchased from Peptides International (Louisville, KY). A23187 was purchased from Calbiochem-Behring (LaJolla, CA). Fluconazole was provided by Pfizer Inc., Groton, CT, disulfonic acid (MOPS), clotrimazole (CLT), miconazole, econazole, metronidazole, and all other drugs and chemicals were

purchased from Sigma Chemical C . (St. Louis, MO) and Fisher Scientific Co. (Fair Lawn, NJ), and the radioisotop 86Rb from Dupont (Billerica, MA)

Assays for Cell Proliferation

DNA. synthesis, assessed by the uptake of [3H]thymidine: Cells are grown in either 48 or 96 wells plates (Costar, Cambridge, MA) at 10⁴ and 0.8 10³ cells per well, respectively, and grown in Dubelcco's modified Eagle's medium (DME, Gibco, Grand Island, NY) supplemented with 10% heat-inactivated calf serum; they are kept at 37°C in 5% CO₂. When they reach confluence, usually between 3 and 4 days, the medium is replaced with DME 0.5% serum to make them quiescent, and mitogenesis assays are performed 24 hours later.

Quiescent cells are exposed to a mitogenic stimulus, such as 10% serum, PDGF (Sigma Co. St. Louis, Mo), bFGF (Upstate Biotechnologies, Lake Placid, NY), or other appropriate mitogen according to the cell line, and 3 hours later 1 µCi/ml of [3H]thymidine (Dupont, Billerica, MA) is added to the wells, and the cells maintained at 37°C/5% CO2 for additional 21 hours. Then the cells are washed 3 times with DME medium and the acid-precipitable radioactivity is extracted with cold 10% TCA (Sigma, Co). After neutralization with 0.3 N NaOH (Sigma Co.), aliquots are counted in a Packard Tri-Carb Scintillation counter (Packard Instrument, Downer's Grove, IL).

Measurement of cell density in culture plates: Cells of a specific test cell line are seeded at precisely the same low density in culture plates and incubated for approximately 12 hours in DME 10% serum, or other culture medium depending on the cell line tested. After 12 hours, the test drug, for example clotrimazole 10 μM , is added to the cell medium of one plate and a similar amount of only the carrier of the drug, for example ethanol 10 μL , to another plate. After 48 to 74 hours, the cell density in control (ethanol) and

experim ntal (clotrimazol) plates is assessed under a light inverted microscope. by measuring the surfac of the culture plate covered by the cell monolayer. Alternatively, the cells can be detached from the plate by incubation with trypsin (Sigma, Co.) 50% (v/v) in ethylene diaminotetraacetic acid (ECTA; Sigma, Co); then the cells are counted in an hemocytometer chamber (Fisher, Pittsburgh, PA).

Assays for Inhibitors of Ca++ Activated K Channel

Ca⁺⁺-sensitive K+ channels have wide distribution among cells, including the human red cell where they were originally discovered and which is the most commonly utilized assay system for activators and inhibitors of the channel for the following reasons: they are readily available, can be easily manipulated in the laboratory, and transport assays can be accurately standardized by reading the hemoglobin concentration of a red cell suspension.

Preparation of Human Red Blood Cells: Blood is collected in heparinized tubes and centrifuged in a Sorvall centrifuge (RB 5B, Du Pont Instruments, Newtown, CT) at 5°C for 10 minutes at 3000 g. Plasma and buffy coat are carefully removed and the cells washed four times with a washing solution containing 150 mM choline chloride (Sigma Co), 1mM MgCl2 (Sigma Co), 10mM Tris-MOPS (Sigma, Ca), pH 7.4 at 4°C(CWS). An aliquot of cells is then suspended in an approximately equal volume of CWS, and from this original cell suspension hematocrit (Hct) and hemoglobin (optical density at 540 nm) are determined.

Methods to Test Inhibitors of the Ca⁺⁺ Activated K: To test inhibitors of the Ca⁺⁺ activated K channel, the channel is activated using the calcium ionophore A23187 (Chalbiochem).

By Atomic Absorption Spectrometry: Washed human erythrocyte are suspended at an hematocrit \simeq 1% in CWS containing 0.150mM CaCl2 (Sigma Co) Aliquots of 1 ml are removed at 0, 3 and 5 minutes, layered on top of 0.3ml of the

oil n-butyl phthalate (Fair Lane, NJ) placed in an Eppendorf microtube (Fisher) and then centrifuged in a micro centrifuge for 20 seconds. At time 5.30 minutes, ionophore A23187 (1µM final concentration) is added and samples removed and spin down through phthalate at times 6, 7, 8 and 9 minutes. The supernatant on top of the oil layer is removed and its K+ concentration is measured by atomic absorption spectrometry using a Perking Elmer model 5000 spectrometer (Perkin Elmer Corp., Norwolk, CT). The efflux of K+ (mmol/l cells/h) in the absence and presence of the inhibitor is calculated from the slope of the curves relating the K+ concentration in the supernatants (mmol/l cells) vs. time (min.).

By radioisotopic measurement of ^{86}Rb influx. The incubation medium is the same but contains 2 mM KCl and 1 $\mu\text{Ci/ml}$ of the radioactive tracer ^{86}Rb . After spinning the samples through the phthalate layer, the tubes are rapidly frozen (-80°C) by immersion in methanol-dry ice, the tips of the tubes containing the packed red cells cut, and counted in a Packard Gamma Counter.

Example 1

The inhibitory effect of clotrimazole (CLT) on cell proliferation was assessed in normal, non-cancerous cells.

Rat vascular smooth muscle cells (murine cell line): Quiescent cells were stimulated with purified growth factors (PDGF and bFGF, 5 μ M) and synthesis of DNA was assessed by the incorporation of [3H]thymidine measured 24 hours later. As shown in Fig. 1, 10 μ M CLT completely inhibited both PDGF and bFGF stimulated DNA synthesis. The effect was not due to a toxic, non-specific, effect because it was reversed by removing CLT and re-stimulating the cells with the corresponding growth factor (Fig. 1).

Example 2

Dose response inhibitors of DNA synthesis by clotrimazole was tested using rat vascular smooth muscle cells as described above. Clotrimazole was tested at concentrations of 0.001 µM, 0.1 µM, 1 µM and 10µM.

Cells were stimulated using 5 µM bFGF. Inhibition was dose dependent, with 45% inhibition at 1 µM and complete inhibition at 10 µM. The ID₅₀ was about 1.5 µM. (Fig. 2)

Example 3

Bovine endothelial (BAEC) and human umbilical vein (HUVEC): Cells were seeded at a low density (2.5 x 10⁵) in cell culture flasks (75 ml flasks) containing DME 10% calf serum (BAEC) or fetal calf serum (HUVEC); after 12 hs, when the cells were attached to the surface of the flasks, CLT (10 µM) or carrier (ethanol) were added to triplicate flasks. After 48 hs cell growth was assessed by optic miscroscopy calculating the surface of the culture flask covered by the cell monolayer. Both BAEC and HUVEC cells had covered 90 ± 2% of the flask surface in the absence and less than 10% in the presence of CLT (data not shown).

Example 4

Other antimycotics were tested for their inhibition of bFGF-stimulated DNA synthesis in rat vascular smooth muscle cells. As shown in Figure 3, at a concentration of 10µM, 3 compounds, CLT, econazole (ECZ) and miconazole (MCZ) inhibited DNA synthesis. The order of inhibitory potency was CLT more potent than ECZ, and ECZ more potent than MCZ. In contrast, other inhibitors of the Ca⁺⁺activated K channel, namely Charybdotoxin, kaliotoxin and iberotoxin, also failed to inhibit DNA synthesis.

Example 5

Th inhibitory ffect of (CLT) on the Ca⁺⁺ activated K channel of human erythrocytes was assessed in the presence of 60 μ mol A23187/L cell and 100 μ MCaCl₂. CLT markedly inhibited the CA⁺⁺ activated 86Rb influx and K efflux. Mean-values of ID₅₀ (calculated with Dixon plot analysis.) was 143 \pm 60 nM(n=3).

Other antimycotics were tested for their inhibition of the Ca++ activated 86Rb influx human erythrocytes. The order of inhibitory potency was clotrimazole more than miconazole; and both of these were more than econazole. There was no inhibition by fluconazole, ornidazole and tinidazole, 2 related compounds, and only marginal with metronidazole a member of the nitroimidazole group (Figure 4).

Example 6

endothelial cells by activated complement. When endothelial cells (EC) in culture (both BAEC and HUVEC) are treated with terminal complement components to form the MAC (membrane attack complex of complement), they release into the culture medium a potent mitogenic activity that stimulates the proliferation of quiescent cells used as indicators of the mitogens. Both, quiescent Swiss 3T3 and vascular smooth muscle cells are stimulated by the mitogens released form EC in-response to the MAC (Figure 5; Halperin et al. unpublished observation). Moreover, immunoprecipitation with specific antibodies has documented that both PDGF and bFGF released from the EC contribute in approximately equal proportion to the mitogenic activity induced by the MAC (data not shown).

To determine whether CLT inhibited the cell proliferative activity released by the MAC from EC, quiescent 3T3 and vascular smooth muscle cells were stimulated in the presence and absence of 10 μM CLT with conditioned media

obtained from MAC treated EC. The results indicate that CLT completely inhibited the proliferative response to mitogene released from EC (Figure 5).

Example 7

The inhibitory effect of clotrimazole (CLT) on cell proliferation was assessed in normal fibroblasts.

Swiss 3T3 cells (murine fibroblast cell line):
Quiescent cells were stimulated with purified growth factors
(PDGF and bFGF, 5 µM) and synthesis of DNA was assessed by
the incorporation of [3H]thymidine measured 24 hours later.
10 µM CLT completely inhibited both PDGF and bFGF
stimulated DNA synthesis. The effect was not due to a toxic,
non-specific, effect because it was reversed by removing CLT
and re-stimulating the cells with the corresponding growth
factor.

Those skilled in the art will be able to ascertain with no more than routine experimentation numerous equivalents to the specific imidazoles and processes described herein. Such equivalents are considered to be within the scope of the invention and are intended to be embraced by the following claims in which we claim:

CLAIMS

1. A method for treating an arteriosclerotic condition, comprising:

administering to a subject in need of such treatment as imidazole that inhibits the Ca⁺⁺ activated potassium channel and that inhibits vascular smooth muscle cell proliferation.

- 2. A method for treating an arteriosclerotic condition as claimed in claim 1 wherein the imidazole is administered to a subject who has sustained an injury to a blood vessel.
- 3. A method for treating an arteriosclerotic condition as claimed in claim 2 wherein the imidazole is administered to a subject who has undergone a balloon angioplasty procedure to prevent restenosis in the subject.
- 4. A method for treating an arteriosclerotic condition as claimed in claim 2 wherein the imidazole is administered to a subject who has undergone vascular surgery.
- 5. A method for treating an arteriosclerotic condition as claimed in claim 1 wherein the imidazole is administered to a subject who has undergone a heart transplant.
- 6. A method for treating an arteriosclerotic condition as claimed in claim 1 wherein the imidazole is administered to a subject who has classical atherosclerosis.
- 7. A method for treating an arteriosclerotic condition as claimed in claim 1 wherein the imidazole is administered to a subject who has diabetes.

- 8. A method for tr ating an art ri sclerotic condition as claimed in any one of claims 1. 2. 3. 4. 5. 6 and 7. wherein clotrimazole is administered to the subject.
- 9. A method for treating an arteriosclerotic condition as claimed in any one of claims 1, 2, 3, 4, 5, 6 and 7, wherein miconazole is administered to the subject.
- 10. A method for treating an arteriosclerotic condition as claimed in any one of claims 1, 2, 3, 4, 5, 6 and 7, wherein econazole is administered to the subject.
- 11. A method for inhibiting vascular smooth muscle cell proliferation comprising contacting vascular smooth muscle cells of a species with an imidazole that inhibits the Ca⁺⁺ activated potassium channel of erythrocytes of the species for the purpose of inhibiting proliferation of said vascular smooth muscle cells.
- 12. A method for inhibiting smooth muscle cell proliferation as claimed in claim 11, wherein the muscle cells are contacted with the imidazole \underline{ex} \underline{vivo} .
- 13. A method for inhibiting smooth muscle cell proliferation as claimed in claim 11 wherein the muscle cells are free of fungus.
- 14. A method as claimed in any one of claims 11, 12 and 13 wherein the smooth muscle cells are contacted with an imidazole selected from the group consisting of: clotrimazole, miconazole and econazole.

PCT/US94/01749

-20-

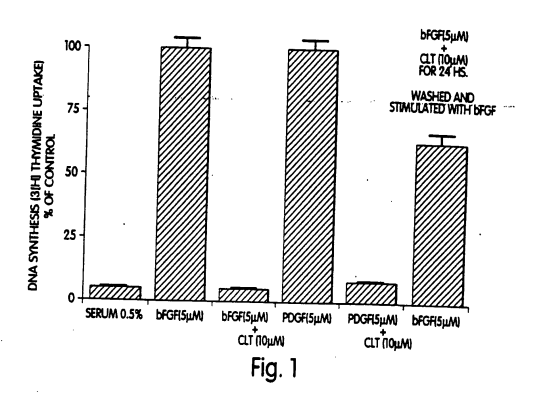
- 15. A method for inhibiting indothelial cell proliferation comprising contacting endoth lial cells of a species with an imidazole that inhibits the Ca⁺⁺ activates potassium channel of erythrocytes of the species for the purpose of inhibiting proliferation of said endothelial cells.
- 16. A method for inhibiting endothelial cells proliferation as claimed in claim 15 wherein the endothelial cells are contracted with the imidazole ex vivo.
- 17. A method for inhibiting endothelial cell proliferation as claimed in claim 15 wherein the endothelial cells are free of fungus.
- 18. A method for inhibiting endothelial cells proliferation as claimed in any one of claims 15, 16 or 17 wherein the endothelial cells are contacted with an imidazole selected from the group consisting of:

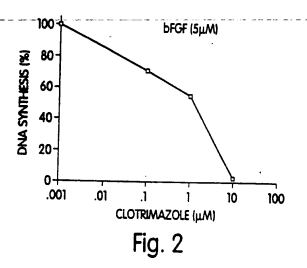
clotrimazole, miconazole and econazole.

- 19. A method for inhibiting fibroblast proliferation comprising contacting fibroblasts of a species with an imidazole that inhibits the Ca⁺⁺ activates potassium channel of erythrocytes of the species for the purpose of inhibiting proliferation of said fibroblasts.
- 20. A method for inhibiting fibroblast proliferation as claimed in claim 19 wherein the fibroblasts are contracted with the imidazole <u>ex vivo</u>.
- 21. A method for inhibiting fibroblast proliferation as claimed in claim 19 wherein the fibroblasts are free of fungus.

22. A method for inhibiting fibroblast prolif ration as claimed in any one of claims 19. 20 or 21 wherein the fibroblasts are contacted with an imidazole selected from the group consisting of:

clotrimazole, miconazole and econazole.





SUBSTITUTE SHEET

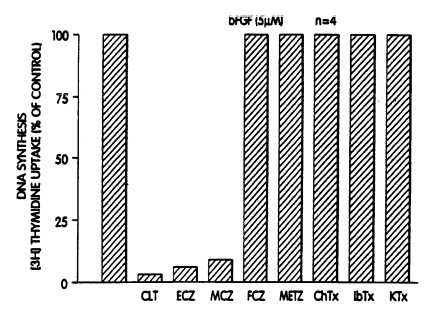
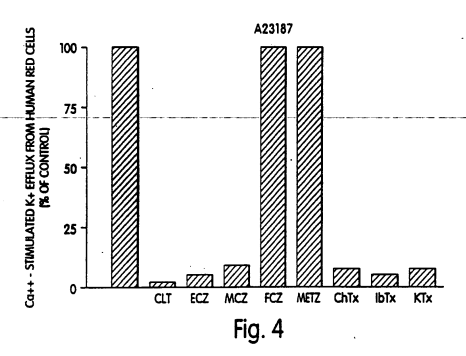


Fig. 3



UBSTITUTE SHEET

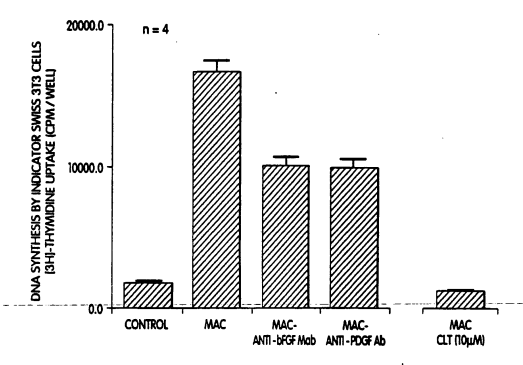


Fig. 5

SUBSTITUTE SHEET

INTERNATIONAL SEARCH REPORT

a. ational application No. PCT/US94/01749

A. CLASSIFICATION OF SUBJECT MATTER						
US CL. 314796.194						
According to International Patent Classification (IPC) or to both national classification and IPC						
	DC CTAPCHED					
Minimum d	locumentation searched (classification system followed	l by classification symbols)				
U.S. :	514/396,824					
Documentar	tion searched other than minimum documentation to the	extent that such documents are included	in the fields searched			
NONE						
	data base consulted during the international search (me					
MUSCLE	iline, embase, medline: econazole, clott :	IIMAZULE, MICUNAZULE,ARI BNO	SCLENUT, SMOUTH			
C. DOC	CUMENTS CONSIDERED TO BE RELEVANT					
Categorya	Citation of document, with indication, where ap	propriets, of the relevant passages	Relevant to claim No.			
	·					
A	US, A, 4,916,118 (Fidler et al.) 10) April 1990.	1-22			
A	US, A, 5,132,315 (Kohn et al.) 21	July 1992.	1-22			
		•				
	·					
i	··		-			
			·			
Further documents are listed in the continuation of Box C. See patent family annex.						
7	notes entegories of clind documents:	"T" hear document published after the let does not not in conflict with the applic	renational filing date or priority when but aims to understand the			
	be part of puriodic relevance	presipts or developing the to "X" document of particular reference the	e chined investor count be			
'E' 4	eller document published on or after the interestional filing data remainst which may throw doubts on priority chain(e) or which is	remittend nevel or expect to execute when the dorument is taken since	ared to ignorive as investive sup-			
ai ai	romant which may throw doubts on priority china(s) or which is sed to combilith the publication data of another chining or other coint roman (so specifies)	"Y" document of posicular relavances to	us oblined invention connect be a step when the document is			
.0	occurrent referring to an oral disclarate, use, exhibition or other	combined with one or more other sec being obvious to a person skilled in t	à decregage, auch combination			
'P' 4	ocument published prior to the International filing data but later than a priority data chained	'A' decement member of the same print				
Date of the actual completion of the international search Date of mailing of the international search report						
30 MARCH 1994 MAY 2 5 1994						
Name and mailing address of the ISA/US Authorized officer						
Commissioner of Petents and Trademarks Box PCT Washington, D.C. 20231 KIMBERLY JORDAN						
	na, D.C. 20231 No. (703) 305-3230	Telephone No. (703) 308-1235	7			

Facsimile No. (703) 305-3230
Form PCT/ISA/210 (second sheet)(July 1992)=